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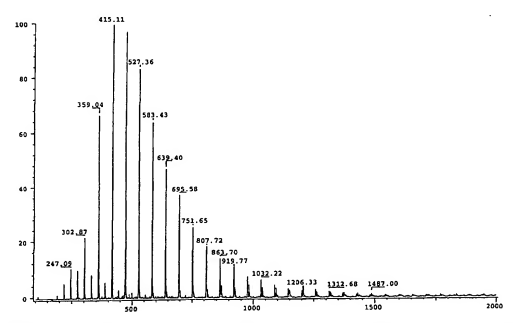
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(54) Title: POLYALKENYL SULFONATES



(57) Abstract: Provided is a polyalkenyl sulfonic acid composition comprising a mixture of polyalkenyl sulfonic acids derived from a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers, and a method for making the composition. Also provided are low and high overbased sulfonates made from the mixture of polyalkenyl sulfonic acids, and lubricating oils containing the sulfonates.



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POLYALKENYL SULFONATES

BACKGROUND OF THE INVENTION

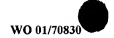
Sulfonates are a class of chemicals used in household, industrial, and institutional cleaning applications, personal care and agricultural products, metalworking fluids, industrial processes, emulsifying agents, corrosion inhibitors and as additives in lubricating oils. Some of the desirable properties of sulfonates for use in lubricating oil applications include their low cost, compatibility, water tolerance, corrosion inhibition, emulsion performance, friction properties, high temperature stability, rust performance, and light color.

Sulfonates that are used in lubricating oil applications have been classified as either neutral sulfonates, low overbased (LOB) sulfonates, or high overbased (HOB) sulfonates.

In the past, natural sulfonates, made as a by-product of white oil and process oil production, dominated the sulfonate market. However, as refineries switched to hydrotreating processes, which gave improved yields of process oils and white oils, and as the desire for higher utilization of raw materials and thus improved economics grew, synthetic sulfonates have become more readily available. Many synthetic sulfonates have been produced from sulfonated polyalkyl aromatic compounds. Unfortunately, many synthetic sulfonates provide properties that are inferior to the properties of the natural sulfonates. Thus, there is a need for low cost synthetic sulfonates that have good performance properties and can serve as a replacement for the natural sulfonates.

SUMMARY OF THE INVENTION

The present invention provides a polyalkenyl sulfonic acid composition comprising a mixture of polyalkenyl sulfonic acids derived from a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers. The present invention also provides such a composition wherein the alkyl vinylidene isomer is a methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer. Also provided is such a composition wherein the number average molecular weight of the polyalkene is about 168 to about 5000. In a preferred embodiment, the polyalkene is polyisobutene. In another preferred embodiment, the



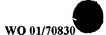
1	polyalkene is polyisobutene and the molecular weight distribution of the
2	polyisobutenyl sulfonic acids has at least 80% of the polyisobutenyl sulfonic acids
3	molecular weights separated by even multiples of 56 daltons. The present invention
4	further provides such a composition wherein the polyalkene is polyisobutene and less
5	than 20% of the polyisobutenyl sulfonic acids in the molecular weight distribution of
6	the polyisobutenyl sulfonic acids contain a total number of carbon atoms that is not
7	evenly divisible by four.
8	Also provided by the present invention is an improved method of making
9	polyalkenyl sulfonic acid by sulfonating polyalkenes, wherein the improvement

Also provided by the present invention is an improved method of making polyalkenyl sulfonic acid by sulfonating polyalkenes, wherein the improvement comprises using as the polyalkenes a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers. The present invention further provides the product of this process.

The present invention further provides a polyalkenyl sulfonate composition having a TBN of about 0 to about 60 wherein the polyalkenyl sulfonate is an alkali metal or alkaline earth metal salt of a polyalkenyl sulfonic acid derived from a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers. Further provided in accordance with this invention is a polyalkenyl sulfonate composition having a TBN of greater than 60 to about 400 wherein the polyalkenyl sulfonate is an alkali metal or alkaline earth metal salt of a polyalkenyl sulfonic acid derived from a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers.

In accordance with the present invention there is also provided an improved method of making polyalkenyl sulfonate by sulfonating polyalkenes and reacting the resulting polyalkenyl sulfonic acid with an alkali metal or alkaline earth metal, the improvement comprising using as the polyalkenes a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers. The present invention also provides the product produced by this process.

Also provided by the present invention is a lubricating oil composition comprising a major amount of an oil of lubricating viscosity and a minor amount of a polyalkenyl sulfonate composition having a TBN of about 0 to about 60 wherein the polyalkenyl sulfonate is an alkali metal or alkaline earth metal salt of a polyalkenyl sulfonic acid derived from a mixture of polyalkenes comprising greater than 20 mole



1	percent alkyl vinylidene and 1,1-dialkyl isomers. The present invention also provides
2	a lubricating oil composition comprising a major amount of an oil of lubricating
3	viscosity and a minor amount of a polyalkenyl sulfonate composition having a TBN of
4	greater than 60 to about 400 wherein the polyalkenyl sulfonate is an alkali metal or
5	alkaline earth metal salt of a polyalkenyl sulfonic acid derived from a mixture of
6	polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl
7	isomers.
8	
9	BRIEF DESCRIPTION OF THE DRAWINGS
10	Figure 1 is a negative ion electrospray ionization mass spectrum of a

Figure 1 is a negative ion electrospray ionization mass spectrum of a polybutene sulfonic acid of a polybutenyl sulfonic acid made in accordance with the present invention.

Figure 2 is a negative ion electrospray ionization mass spectrum of a polybutene sulfonic acid made from a polybutene with less than 10% methylvinylidene isomer content, i.e., not a polybutenyl sulfonic acid of this invention.

DETAILED DESCRIPTION OF THE INVENTION

The polyalkenyl sulfonic acids of this invention are prepared by reacting a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers with a source of sulfur trioxide -SO₃-. The source of -SO₃- can be a mixture of sulfur trioxide and air, sulfur trioxide hydrates, sulfur trioxide amine complexes, sulfur trioxide ether complexes, sulfur trioxide phosphate complexes, acetyl sulfate, a mixture of sulfur trioxide and acetic acid, sulfamic acid, alkyl sulfates or chlorosulfonic acid. The reaction may be conducted neat or in any inert anhydrous solvent. The conditions for sulfonation are not critical. Reaction temperatures can range from -30°C. to 200°C. and depends on the particular sulfonating agent employed. For example, acetyl sulfate requires low temperatures for reaction and elevated temperatures should be avoided to prevent decomposition of the product. Reaction time can vary from a few minutes to several hours depending on other conditions, such as reaction temperature. The extent of the reaction can be determined

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1 by titration of 2 Typical mole 3 The p. 4 and acetic and

- by titration of sulfonated polyalkene after any free sulfuric acid has been washed out.
- 2 Typical mole ratios of sulfonating agent to polyalkene can be about 1:1 to 2:1.

The preferred sulfonating agent is acetyl sulfate (or a mixture of sulfuric acid and acetic anhydride which forms acetyl sulfate *in situ*) which produces the polyalkenyl sulfonic acid directly. Other sulfonating agents, such as a mixture of sulfur trioxide and air, may produce a sultone intermediate that needs to be hydrolyzed to the sulfonic acid. This hydrolysis step can be very slow.

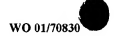
The polyalkenes used to prepare the polyalkenyl sulfonic acid are a mixture of polyalkenes having 12 to 350 carbon atoms. The mixture comprises greater than 20 mole percent, preferably greater than 50 mole percent, and more preferably greater than 70 mole percent alkylvinylidene and 1,1-dialkyl isomers. The preferred alkylvinylidene isomer is a methyl vinylidene isomer, and the preferred 1,1-dialkyl isomer is a 1,1-dimethyl isomer.

The polyalkenes have a number average molecular weight in the range of about 168 to about 5000. Polyalkenes having number average molecular weights of about 550, 1000 or 2300 are particularly useful.

The preferred polyalkene is polyisobutene. Especially preferred are polyisobutenes made using BF₃ as catalyst.

U. S. Patent No. 5,408,018, which issued on April 18, 1995 to Rath and which is incorporated by reference in its entirety, and the references cited therein describe a suitable process for the production of polyisobutenes that contain greater than 20 mole percent alkylvinylidene and 1,1-dialkyl isomers.

Typically, when polyisobutenyl sulfonic acids or sulfonates are prepared from polyisobutene having a low mole percent of alkylvinylidene and 1,1-dialkyl isomers, the product has a molecular weight distribution similar to that shown in Figure 2. Since polyisobutene is used to prepare the sulfonic acid or sulfonate, it should be expected that the mass spectrum of the product would show compounds separated by even multiples of 56 daltons, i.e., a C₄H₈ fragment. However, Figure 2, which is the mass spectrum of a polyisobutenyl sulfonate prepared from a polyisobutene having a mole percent of methylvinylidene isomers of less than 20%, clearly shows compounds which are separated by less than 56 daltons.

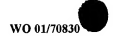


It has now been discovered that when polyisobutene having a mole percent of alkyl vinylidene and 1,1-dialkyl isomers greater than 20% is used to prepare polyisobutenyl sulfonic acids or sulfonates, the molecular weight distribution of the resulting product has at least 80% of the polyisobutenyl sulfonic acids or sulfonates whose molecular weights are separated by even multiples of 56 daltons (see Figure 1). In other words, less than 20% of the polyisobutenyl sulfonic acids or sulfonates in the molecular weight distribution of the sulfonic acids or sulfonates contain a total number of carbon atoms that is not evenly divisible by four.

The polyalkenyl sulfonates of this invention are prepared by reacting the polyalkenyl sulfonic acid (prepared as described above) with a source of an alkali or alkaline earth metal. The alkali or alkaline earth metal can be introduced into the sulfonate by any suitable means. One method comprises combining a basically reacting compound of the metal, such as the hydroxide, with the polyalkenyl sulfonic acid. This is generally carried out in the presence of a hydroxylic promoter such as water, alcohols such as 2-ethyl hexanol, methanol or ethylene glycol, and an inert solvent for the sulfonate, typically with heating. Under these conditions, the basically reacting compound will yield the metal sulfonate. The hydroxylic promoter and solvent can then be removed to yield the metal sulfonate.

Under certain circumstances, it may be more convenient to prepare an alkali metal polyalkenyl sulfonate and convert this material by metathesis into an alkaline earth metal sulfonate. Using this method, the sulfonic acid is combined with a basic alkali metal compound such as sodium or potassium hydroxide. The sodium or potassium sulfonate obtained can be purified by aqueous extraction. Then, the sodium or potassium sulfonate is combined with an alkaline earth metal salt to form the alkaline earth metal sulfonate. The most commonly used alkaline earth metal compound is a halide, particularly a chloride. Typically, the sodium or potassium sulfonate is combined with an aqueous chloride solution of the alkaline earth metal and stirred for a time sufficient for metathesis to occur. Thereafter, the water phase is removed and the solvent may be evaporated, if desired.

The preferred sulfonates are alkaline earth metal sulfonates, especially those of calcium, barium and magnesium. Most preferred are the calcium and magnesium sulfonates.



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The polyalkenyl sulfonates of this invention are either neutral or overbased sulfonates. Overbased materials are characterized by a metal content in excess of that which would be present according to the stoichiometry of the metal cation in the sulfonate said to be overbased. Thus, a monosulfonic acid when neutralized with an alkaline earth metal compound, such as a calcium compound, will produce a normal sulfonate containing one equivalent of calcium for each equivalent of acid. In other words, the normal metal sulfonate will contain one mole of calcium for each two moles of the monosulfonic acid.

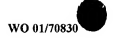
By using well known procedures, overbased or basic complexes of the sulfonic acid can be obtained. These overbased materials contain amounts of metal in excess of that required to neutralize the sulfonic acid. Highly overbased sulfonates can be prepared by the reaction of overbased sulfonates with carbon dioxide under reaction conditions. A discussion of the general methods for preparing overbased sulfonates and other overbased products is disclosed in U.S. Patent No. 3,496,105, issued February 17, 1970 to LeSuer, which in incorporated by reference in its entirety.

The amount of overbasing can be expressed as a Total Base Number ("TBN"), which refers to the amount of base equivalent to one milligram of KOH in one gram of sulfonate. Thus, higher TBN numbers reflect more alkaline products and therefor a greater alkalinity reserve. The TBN for a composition is readily determined by ASTM test method D664 or other equivalent methods. The overbased polyalkenyl sulfonates of this invention can have relatively low TBN, i.e., about 0 to about 60, or relatively high TBN, i.e., greater than 60 to about 400.

The polyalkenyl sulfonates of this invention are useful as additives in lubricating oils. They have good tolerance to water, a light color and provide good performance characteristics.

The lubricating oil compositions of this invention comprise a major amount of an oil of lubricating viscosity and a minor amount of the polyalkenyl sulfonates of this invention. The oils can be derived from petroleum or be synthetic. The oils can be paraffinic, naphthenic, halosubstituted hydrocarbons, synthetic esters, or combinations thereof. Oils of lubricating viscosity have viscosities in the range from 35 to 55,000 SUS at 100°F., and more usually from about 50 to 10,000 SUS at 100°F. The

32 lubricating oil compositions contain an amount of the polyalkenyl sulfonates of this



1	invention sufficient to provide dispersant properties, typically from about 0.1 weight
2	percent to 10 weight percent, preferably from about 0.5 weight percent to about 7
3	weight percent.
4	Other conventional additives that can be used in combination with the
5	polyalkenyl sulfonates of this invention include oxidation inhibitors, antifoam agents,
6	viscosity index improvers, pour point depressants, dispersants and the like.
7	The lubricating oil compositions of this invention are useful for lubricating
8	internal combustion engines and automatic transmissions, and as industrial oils such
9	as hydraulic oils, heat transfer oils, torque fluids, etc.
10	
11	EXAMPLE 1
12	PREPARATION OF A POLYISOBUTENE SULFONIC ACID FROM A
13	HIGH METHYLVINYLIDENE POLYISOBUTENE THAT HAS
14	A Mn OF 550 AND ACETYL SULFATE
15	To a beaker is added 5.5g (0.01 mol) of Glissopal 550 polyisobutene (which
16	has greater than about 80% methylvinylidene content with a number average
17	molecular weight of about 550) dissolved in 20mL hexane. To this is added 1.63g
18	acetic anhydride (0.016 mole) and then 0.98g sulfuric acid (0.01 mole). The resulting
19	mixture is stirred at room temperature for one hour. Then some methanol is added to
20	quench the reaction and the solvents are removed in vacuo. A total of 7.16g of crude
21	polyisobutene sulfonic acid is obtained.
22	EXAMPLE 2
23	PREPARATION OF A LOB POLYISOBUTENE SODIUM SULFONATE
24	FROM THE PRODUCT OF EXAMPLE 1
25	To 5.91g of the sulfonic acid from Example 1 is added 20 mL isopropyl
26	alcohol and 1 g sodium hydroxide in 1 mL of water. The resulting mixture is refluxed
27	for 6 hours and then held at room at room temperature overnight. Two layers form
28	and the bottom layer is decanted. The top layer contains sodium polyisobutene
29	sulfonate (5.67g) which contains 88% actives. The lower layer is stripped in vacuo
30	and contains 1.14g of a mixture of sodium hydroxide and sodium polyisobutene
31	sulfonate.



1	EXAMPLE 3
2	PREPARATION OF CALCIUM LOB POLYISOBUTENE SULFONATE
3	To a 2L round bottom flask is added 500g of Glissopal 550 polyisobutene
4	(0.91 mol), 140.3g acetic anhydride (1.38 mol), and 84.7g concentrated sulfuric acid
5	(0.86 mol) at room temperature. The resulting mixture is stirred 4 hours at room
6	temperature. Then to this mixture is added 50 mL methanol to quench the reaction,
7	and 500g of 100 neutral diluent oil. To this is then added 32.0g calcium hydroxide
8	(0.43 mol) and 20 mL water. The resulting mixture is heated to 175°F and then 100
9	mL water is added. This is then heated to 225-230°F for 30 minutes, and then heated
10	at 330°F for 1 hour to strip off the water. A calcium polyisobutene sulfonate is
11	obtained.
12	EXAMPLE 4
13	PREPARATION OF POLYISOBUTENE SULFONIC ACID USING SO3 AND AIR
14	A thin film of Glissopal 550 polyisobutene is sulfonated using SO ₃ and air
15	under the following conditions: temperature 60°C, SO ₃ flow 16L/hr, air flow 192L/hr
16	feed rate 4.5g/min. The product from this reaction is a mixture of polyisobutene
17	sulfonic acid and polyisobutene sultone. The product contains 2.04% sulfonate as
18	calcium sulfonate and 0.70% sulfuric acid as determined by hyamine titration.
19	EXAMPLE 5
20	PREPARATION OF POLYISOBUTENE SULFONIC ACID USING SO3 AND AIR
21	A thin film of Glissopal 550 polyisobutene is sulfonated using SO ₃ and air
22	under the following conditions: temperature 60°, SO ₃ flow 16L/hr, air flow 192L/hr,
23	feed rate 4.2g/min. A total of 1354g product is obtained which is a mixture of
24	polyisobutene sulfonic acid and polyisobutene sultone. The product contains 2.5%
25	sulfonate as calcium sulfonate and 1.02% sulfuric acid as determined by hyamine
26	titration. The acid number is determined by the ASTM D664 test to be 59.9 mg
27	KOH/g sample.
28	EXAMPLE 6
29	PREPARATION OF SODIUM POLYISOBUTENE SULFONATE
30	The mixture of polyisobutene sulfonic acid and polyisobutene sultone from
31	Example 5 is hydrolyzed using the following procedure. To a 100 mL three neck
32	flask equipped with a reflux condenser and stirrer is added 20g of polyisobutene



1	suitonic acid and the resulting mixture is heated to 100°C. To this is added 5 mL 49%			
2	sodium hydroxide solution and the resulting mixture is stirred for four hours. The			
3	product from this reaction is a mixture of sodium polyisobutene sulfonate and			
4	polyisobutene sultone.			
5	EXAMPLE 7			
6	PREPARATION OF CALCIUM POLYISOBUTENE SULFONATE			
7	FROM 550 MW POLYISOBUTENE (NEUTRAL SULFONATE)			
8	To a 2 L round bottom flask is added 500g (0.91 mol) Glissopal 550 (550 M_n			
9	polyisobutene containing about 85% methylvinylidene isomer), 140.3 g acetic			
10	anhydride, (1.38 mol; 1.5 equivalents), and 84.7g sulfuric acid (0.864 mol; 0.95			
11	equivalents) dropwise at room temperature. The resulting mixture is stirred 4 hours			
12	at room temperature. Then to this is added 50 mL methanol and then 500g 100			
13	neutral diluent oil is added. To this is then added 32.0g calcium hydroxide (0.43 mol)			
14	and 20 mL water. This is heated to about 80°C and an additional 100 mL or water is			
15	added. Then the volatile materials are removed at elevated temperatures. The produc			
16	is filtered to give 842.3 g of product which has a TBN of 3.4 mg KOH/g sample, a			
17	viscosity @100°C of 72.4 cSt,, 1.50% Ca, and 2.31% S.			
18	EXAMPLE 8			
19	PREPARATION OF CALCIUM POLYISOBUTENE SULFONATE			
20	FROM 550 MW POLYISOBUTENE (LOB SULFONATE)			
21	To a 4 L beaker is added 500g (0.91 mol) Glissopal 550 polyisobutene (550			
22	M _n polyisobutene with about 85% methylvinylidene isomer content), 140.3 g acetic			
23	anhydride (1.38 mol), and 84.7g sulfuric acid (0.864 mol). The resulting mixture is			
24	stirred 1 hour at room temperature. To this is then added 50 mL methanol, 500g 100			
25	neutral diluent oil, and 100 mL water. The resulting mixture is heated to 190°F and			
26	48g (0.649 mol) calcium hydroxide is added. This is stirred for one hour and then the			
27	temperature is raised to 212°F and maintained there until all the volatile material has			
28	distilled. The resulting product is then filtered to give a LOB calcium polyisobutene			
29	sulfonate which has a TBN of 12.3 mg KOH/g sample, 2.24 % S, 1.85% Ca, and a			
30	viscosity @ 100°C of 79.4 cSt.			



EXAMPLE 9

2 PREPARATION OF ADDITIONAL POLYISOBUTENE SULFONATES

Additional examples of calcium polyisobutene sulfonates are carried out using 4 different conditions, charge mole ratios ("CMR's") and polyisobutene ("PIB") molecular weights as shown in Table 1.

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Ex.	PIB	%	Ac ₂ O	H ₂ SO ₄ /	H ₂ SO ₄ /	%	%	TBN	Vis @
1	M_n	diluent	/	PIB	Ca(OH) ₂	Ca	S		100°C
		oil	PIB						
7	550	50	1.52	0.95	2.01	1.50	2.31	3.4	72.4
8	550	50	1.52	0.95	1.33	1.85	2.24	12.0	79.4
9	550	45	1.51	0.95	1.83	1.84	2.60	5.5	113.8
10	550	45	1.51	0.95	2.00	1.71	2.58	2.6	167.4
11	1000	45	1.51	0.95	1.81	1.08	1.50	3.1	153.8
12	1000	45	1.51	0.95	1.76	1.03	1.52	0.9	156.1
13	1000	45	1.50	0.95	1.83	1.08	1.49	3.9	163.8

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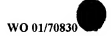
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EXAMPLE 14

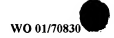
PREPARATION OF CALCIUM ACETATE-FREE 550 Mn

CALCIUM POLYISOBUTENE SULFONATE

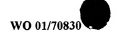
The sulfonic acid from 550 M_n polyisobutene is first prepared by reacting Glissopal 550 polyisobutene (2000 g, 3.64 mol), with 408.3 g acetic anhydride (4.0 mol), and 338.7 g sulfuric acid (3.46 mol). The resulting mixture is stirred for one hour at room temperature. Then 200 mL of methanol is added. The resulting product contains about 90% actives. Then 260 g of this product is diluted with 260 g of 100 neutral diluent oil and this is heated at 40°C with a nitrogen sparge to remove the unreacted acetic acid, methyl acetate, and methanol. Analysis by 'H NMR spectroscopy indicates that only about 0.3% acetic acid remained. This product (448.2 g) is then placed in a 1000 mL beaker and heated to 190°F and to this is added 13.2g calcium hydroxide. The resulting mixture is stirred for 1 hour at 190°F, and then the temperature is increased to 330°F to remove any volatile material. The resulting product is then filtered to give a calcium acetate-free calcium polyisobutene



1	sulfonate which has a TBN of 5.4 mg KOH/g sample, 1.12%Ca, 1.82% S, and a
2	viscosity @ 100°C of 27.5 cSt.
3	COMPARATIVE EXAMPLE A
4	PREPARATION OF 950 Mn POLYISOBUTENE SULFONIC ACID
5	FROM PARAPOL 1000
6	200g Parapol 950 (950 Mn polyisobutene with less than 5% methylvinylidene
7	isomer content, 0.21 mol) is reacted with 22.46g acetic anhydride (0.22 mol) and
8	18.63g sulfuric acid (0.190 mol). The resulting product is stirred at room temperature
9	for 1 hour then 20 mL methanol was added. This product contains only about 67%
10	actives.
11	EXAMPLE 15
12	PREPARATION OF 1000 M_n POLYISOBUTENE SULFONIC ACID
13	FROM GLISSOPAL 1000
14	To 2000g (2.0 mol) of Glissopal 1000 polyisobutene (M_n 1000 with about
15	85% methylvinylidene isomer content) is added 224.6 g acetic anhydride (2.2 mol)
16	and 186.3 g of sulfuric acid (1.90 mol). The resulting product is reacted as in
17	Comparative Example A. The product contains about 90% actives. This shows the
8	improvement in yield that is obtained with the teachings of this invention.
9	COMPARATIVE EXAMPLE B
20	ELECTROSPRAY IONIZATION-MASS SPECTRUM OF SULFONIC ACID
21	FROM POLYISOBUTENE WITH LESS THAN 20% METHYLVINYLIDENE
22.	CONTENT
23	Figure 2 shows the electrospray ionization mass spectrum of a polybutene
24	sulfonic acid from Hivis 5 (polybutene with less than 10% methylvinylidene isomer
25	content). The spectrum shows a molecular weight distribution with molecular ions
26	that are separated by 14 daltons. This indicates that the polyisobutene sulfonic acid
27	actually is not a mixture of C ₁₂ , C ₁₆ , C ₂₀ etc. isomers, but is a mixture of C ₁₂ , C ₁₃ , C ₁₄ ,
28	etc., isomers.



1	EXAMPLE 16
2	ELECTROSPRAY IONIZATION-MASS SPECTRUM OF SULFONIC ACID
3	FROM POLYISOBUTENE WITH MORE THAN 20% METHYLVINYLIDENE
4	CONTENT
5	Figure 1 shows the electrospray ionization mass spectrum of a polybutene
6	sulfonic acid from Glissopal 550 (polybutene with greater than 85% methylvinylidene
7	isomer content). The spectrum shows a molecular weight distribution with molecular
8	ions that are separated by 56 daltons. This indicates that the polyisobutene sulfonic
9	acid is a mixture of C ₁₂ , C ₁₆ , C ₂₀ , etc., isomers (i.e., the ions are multiples of four
10	carbon atoms).



1 WHAT IS CLAIMED IS:

2

A polyalkenyl sulfonic acid composition comprising a mixture of polyalkenyl sulfonic acids derived from a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl isomers.

6

7 2. The composition of claim 1 wherein the mixture of polyalkenes comprises 8 greater than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.

9

The composition of claim 1 wherein the mixture of polyalkenes comprises greater than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.

12

The composition of claim 1, 2 or 3 wherein the alkyl vinylidene isomer is a methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.

15

16 5. The composition of claim 1 wherein the number average molecular weight of 17 the polyalkene is about 168 to about 5000.

18

19 6. The composition of claim 1 wherein the number average molecular weight of 20 the polyalkene is about 550.

21

The composition of claim 1 wherein the number average molecular weight of the polyalkene is about 1000.

2425

8. The composition of claim 1 wherein the number average molecular weight of the polyalkene is about 2300.

27

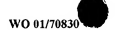
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28 9. The composition of claim 1 wherein the polyalkene is polyisobutene.

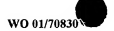
29

The composition of claim 9 wherein the polyisobutene is made using a BF₃
 catalyst.

32

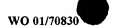


I	11.	The composition of claim I wherein the polyalkene is polyisobutene and the
2		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
3		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
4		multiples of 56 daltons.
5		
6	12.	The composition of claim 1 wherein the polyalkene is polyisobutene and less
7		than 20% of the polyisobutenyl sulfonic acids in the molecular weight
8		distribution of the polyisobutenyl sulfonic acids contain a total number of
9		carbon atoms that is not evenly divisible by four.
10		
l 1	13.	In a method of making polyalkenyl sulfonic acid by sulfonating polyalkenes,
12		the improvement comprising using as the polyalkenes a mixture of polyalkenes
13		comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl
14		isomers.
5		
6	14.	The method of claim 13 wherein the mixture of polyalkenes comprises greater
7		than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
8		
9	15.	The method of claim 13 wherein the mixture of polyalkenes comprises greater
20		than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
21		
22	16.	The method of claim 13, 14 or 15 wherein the alkyl vinylidene isomer is a
23		methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.
24		
25	17.	The method of claim 13 wherein the number average molecular weight of the
26		polyalkene is about 168 to about 5000.
27		
28	18.	The method of claim 13 wherein the number average molecular weight of the
9		polyalkene is about 550.
0		
1	19.	The method of claim 13 wherein the number average molecular weight of the
2		polyalkene is about 1000.



1	20.	The method of claim 13 wherein the number average molecular weight of the
2		polyalkene is about 2300.
3		
4	21.	The method of claim 13 wherein the polyalkene is polyisobutene.
5		
6	22.	The method of claim 21 wherein the polyisobutene is made using a BF ₃
7		catalyst.
8		
9	23.	The method of claim 13 wherein the polyalkene is polyisobutene and the
10		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
11		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
12		multiples of 56 daltons.
13		
14	24.	The method of claim 13 wherein the polyalkene is polyisobutene and less than
15		20% of the polyisobutenyl sulfonic acids in the molecular weight distribution
16		of the polyisobutenyl sulfonic acids contain a total number of carbon atoms
17		that is not evenly divisible by four.
18		
19	25.	The product produced by the method of claim 13, 14, 15, 17, 18, 19, 20, 21,
20		22, 23 or 24.
21		
22	26.	The product produced by the method of claim 16.
23		
24	27.	A polyalkenyl sulfonate composition having a TBN of about 0 to about 60
25		wherein the polyalkenyl sulfonate is an alkali metal or alkaline earth metal salt
26		of a polyalkenyl sulfonic acid derived from a mixture of polyalkenes
27		comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl
28		isomers.
29		
30	28.	The composition of claim 27 wherein the mixture of polyalkenes comprises
31		greater than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
32		

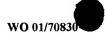
1	29.	The composition of claim 27 wherein the mixture of polyalkenes comprises
2		greater than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
3		
4	30.	The composition of claim 27, 28 or 29 wherein the alkyl vinylidene isomer is a
5		methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.
6		
7	31.	The composition of claim 27 wherein the number average molecular weight of
8		the polyalkene is about 168 to about 5000.
9		
10	32.	The composition of claim 27 wherein the number average molecular weight of
11		the polyalkene is about 550.
12		
13	33.	The composition of claim 27 wherein the number average molecular weight of
14		the polyalkene is about 1000.
15		
16	34.	The composition of claim 27 wherein the number average molecular weight of
17		the polyalkene is about 2300.
18		
19	35.	The composition of claim 27 wherein the polyalkene is polyisobutene.
20		
21	36.	The composition of claim 35 wherein the polyisobutene is made using a BF ₃
22		catalyst.
23		
24	37.	The composition of claim 27 wherein the polyalkene is polyisobutene and the
25		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
26		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
27		multiples of 56 daltons.
28		
29	38.	The composition of claim 27 wherein the polyalkene is polyisobutene and less
30		than 20% of the polyisobutenyl sulfonic acids in the molecular weight
31		distribution of the polyisobutenyl sulfonic acids contain a total number of
32		carbon atoms that is not evenly divisible by four.



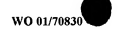
1	39.	A polyalkenyl sulfonate composition having a TBN of greater than 60 to about
2		400 wherein the polyalkenyl sulfonate is an alkali metal or alkaline earth metal
3		salt of a polyalkenyl sulfonic acid derived from a mixture of polyalkenes
4		comprising greater than 20 mole percent alkyl vinylidene and 1,1-dialkyl
5		isomers.
6		
7	40.	The composition of claim 39 wherein the mixture of polyalkenes comprises
8		greater than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
9		
10	41.	The composition of claim 39 wherein the mixture of polyalkenes comprises
11		greater than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
12		
13	42.	The composition of claim 39, 40 or 41 wherein the alkyl vinylidene isomer is a
14		methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.
15		
16	43.	The composition of claim 39 wherein the number average molecular weight of
17		the polyalkene is about 168 to about 5000.
18		
19	44.	The composition of claim 39 wherein the number average molecular weight of
20		the polyalkene is about 550.
21		
22	45.	The composition of claim 39 wherein the number average molecular weight of
23		the polyalkene is about 1000.
24		
25	46.	The composition of claim 39 wherein the number average molecular weight of
26		the polyalkene is about 2300.
27		
28	47.	The composition of claim 39 wherein the polyalkene is polyisobutene.
29		
30	48.	The composition of claim 47 wherein the polyisobutene is made using a BF ₃
31		catalyst.
32		



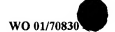
I	49.	The composition of claim 39 wherein the polyaikene is polyisobulene and the
2		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
3		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
4		multiples of 56 daltons.
5		
6	50.	The composition of claim 39 wherein the polyalkene is polyisobutene and less
7		than 20% of the polyisobutenyl sulfonic acids in the molecular weight
8		distribution of the polyisobutenyl sulfonic acids contain a total number of
9		carbon atoms that is not evenly divisible by four.
10		
11	51.	In a method of making polyalkenyl sulfonate by sulfonating polyalkenes and
12		reacting the resulting polyalkenyl sulfonic acid with an alkali metal or alkaline
13		earth metal, the improvement comprising using as the polyalkenes a mixture of
14		polyalkenes comprising greater than 20 mole percent alkyl vinylidene and
15		1,1-dialkyl isomers.
16		
17	52.	The method of claim 51 wherein the mixture of polyalkenes comprises greater
18		than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
19		
20	53.	The method of claim 51 wherein the mixture of polyalkenes comprises greater
21		than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
22		
23	54.	The method of claim 51, 52 or 53 wherein the alkyl vinylidene isomer is a
24		methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.
25		
26	55.	The method of claim 51 wherein the number average molecular weight of the
27		polyalkene is about 168 to about 5000.
28		
29	56.	The method of claim 51 wherein the number average molecular weight of the
30		polyalkene is about 550.
31		



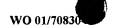
i	57.	The method of claim 31 wherein the number average molecular weight of the
2		polyalkene is about 1000.
3		
4	58.	The method of claim 51 wherein the number average molecular weight of the
5		polyalkene is about 2300.
6		
7	59.	The method of claim 51 wherein the polyalkene is polyisobutene.
8		
9	60.	The method of claim 59 wherein the polyisobutene is made using a BF ₃
10		catalyst.
11		
12	61.	The method of claim 51 wherein the polyalkene is polyisobutene and the
13		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
14		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
15		multiples of 56 daltons.
16		
17	62.	The method of claim 51 wherein the polyalkene is polyisobutene and less than
18		20% of the polyisobutenyl sulfonic acids in the molecular weight distribution
19		of the polyisobutenyl sulfonic acids contain a total number of carbon atoms
20		that is not evenly divisible by four.
21		
22	63.	The product produced by the method of claim 51, 52, 53, 55, 56, 57, 58, 59,
23		60, 61 or 62.
24		
25	64.	The product produced by the method of claim 54.
26		
27	65.	A lubricating oil composition comprising a major amount of an oil of
28		lubricating viscosity and a minor amount of a polyalkenyl sulfonate
29		composition having a TBN of about 0 to about 60 wherein the polyalkenyl
30		sulfonate is an alkali metal or alkaline earth metal salt of a polyalkenyl
31		sulfonic acid derived from a mixture of polyalkenes comprising greater than
32		20 mole percent alkyl vinylidene and 1,1-dialkyl isomers.



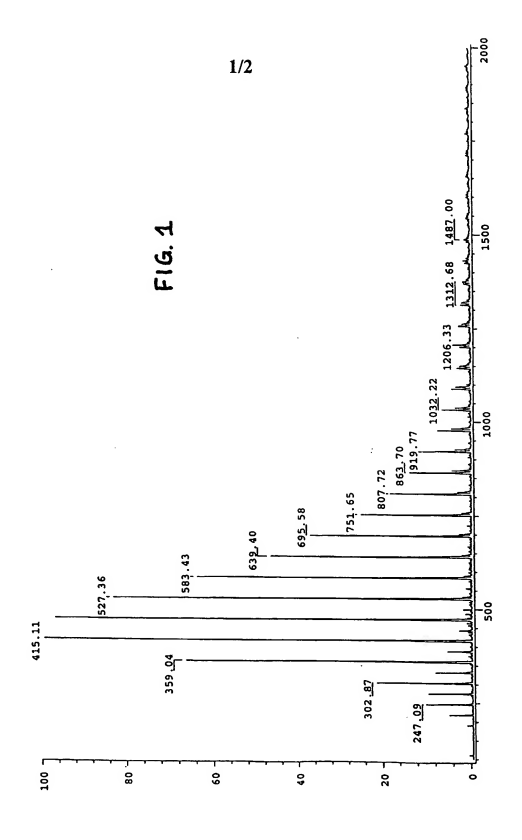
1	66.	The composition of claim 65 wherein the mixture of polyalkenes comprises
2		greater than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
3		
4	67.	The composition of claim 65 wherein the mixture of polyalkenes comprises
5		greater than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
6		
7	68.	The composition of claim 65, 66 or 67 wherein the alkyl vinylidene isomer is a
8		methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.
9		
10	69.	The composition of claim 65 wherein the number average molecular weight of
11		the polyalkene is about 168 to about 5000.
12		
13	70.	The composition of claim 65 wherein the number average molecular weight of
14		the polyalkene is about 550.
15		
16	71.	The composition of claim 65 wherein the number average molecular weight of
17		the polyalkene is about 1000.
18		
19	72.	The composition of claim 65 wherein the number average molecular weight of
20		the polyalkene is about 2300.
21		
22	73.	The composition of claim 65 wherein the polyalkene is polyisobutene.
23		
24	74.	The composition of claim 73 wherein the polyisobutene is made using a BF ₃
25		catalyst.
26		
27	75 .	The composition of claim 65 wherein the polyalkene is polyisobutene and the
28		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
29		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
30		multiples of 56 daltons.
31		

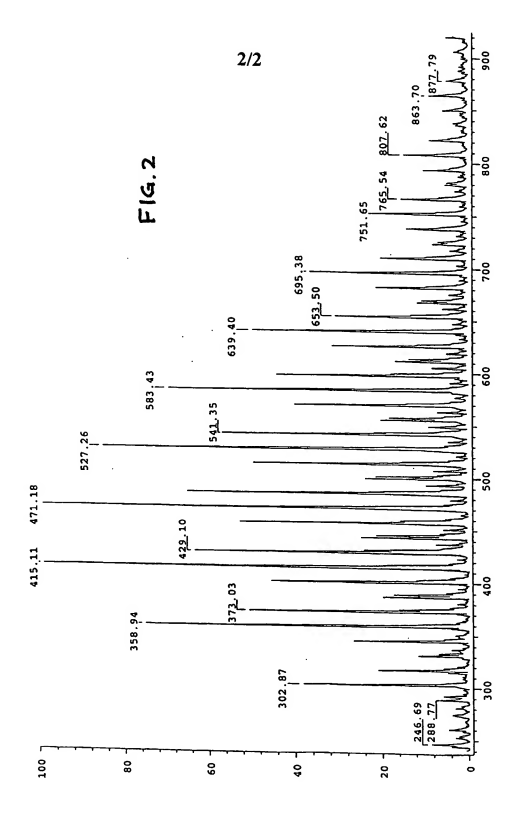


1	76.	The composition of claim 65 wherein the polyalkene is polyisobutene and less
2		than 20% of the polyisobutenyl sulfonic acids in the molecular weight
3		distribution of the polyisobutenyl sulfonic acids contain a total number of
4		carbon atoms that is not evenly divisible by four.
5		
6	77.	A composition comprising a major amount of an oil of lubricating viscosity
7		and a minor amount of a polyalkenyl sulfonate composition having a TBN of
8		greater than 60 to about 400 wherein the polyalkenyl sulfonate is an alkali
9		metal or alkaline earth metal salt of a polyalkenyl sulfonic acid derived from a
10		mixture of polyalkenes comprising greater than 20 mole percent alkyl
1		vinylidene and 1,1-dialkyl isomers.
12		
13	78.	The composition of claim 77 wherein the mixture of polyalkenes comprises
14		greater than 50 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
15		
6	79.	The composition of claim 77 wherein the mixture of polyalkenes comprises
17		greater than 70 mole percent alkyl vinylidene and 1,1-dialkyl isomers.
8		
9	80.	The composition of claim 75, 76 or 77 wherein the alkyl vinylidene isomer is a
20		methyl vinylidene isomer, and the 1,1-dialkyl isomer is a 1,1-dimethyl isomer.
21		
22	81.	The composition of claim 77 wherein the number average molecular weight of
23		the polyalkene is about 168 to about 5000.
24		
25	82.	The composition of claim 77 wherein the number average molecular weight of
26	•	the polyalkene is about 550.
27		
28	83.	The composition of claim 77 wherein the number average molecular weight of
29		the polyalkene is about 1000.
0		•
31	84.	The composition of claim 77 wherein the number average molecular weight of
2		the polyalkene is about 2300.



1	85.	The composition of claim 77 wherein the polyalkene is polyisobutene.
2		
3	86.	The composition of claim 85 wherein the polyisobutene is made using a BF ₃
4		catalyst.
5		
6	87.	The composition of claim 77 wherein the polyalkene is polyisobutene and the
7		molecular weight distribution of the polyisobutenyl sulfonic acids has at least
8		80% of the polyisobutenyl sulfonic acids molecular weights separated by even
9		multiples of 56 daltons.
10		
11	88.	The composition of claim 77 wherein the polyalkene is polyisobutene and less
12		than 20% of the polyisobutenyl sulfonic acids in the molecular weight
13		distribution of the polyisobutenyl sulfonic acids contain a total number of
14		carbon atoms that is not evenly divisible by four.









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- (25) Filing Language:

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09/527,166

17 March 2000 (17.03.2000) US

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- (74) Agents: STUMPF, Walter, L. et al.; Chevron Corporation. Law Dept., P.O. Box 6006, San Ramon, CA 94583-0806 (US).

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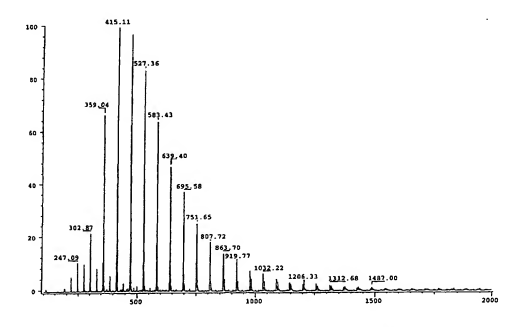
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[Continued on next page]

(54) Title: POLYALKENYL SULFONATES



(57) Abstract: Provided is a polyalkenyl sulfonic acid composition comprising a mixture of polyalkenyl sulfonic acids derived from a mixture of polyalkenes comprising greater than 20 mole percent alkyl vinylidene and 1.1-dialkyl isomers, and a method for making the composition. Also provided are low and high overbased sulfonates made from the mixture of polyalkenyl sulfonic acids, and lubricating oils containing the sulfonates.



01/70830 A3





For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.



al Application No PCT/US 01/08345

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C08F8/36 C08F8/44

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

 $\begin{array}{lll} \mbox{Minimum documentation searched (classification system followed by classification symbols)} \\ \mbox{IPC 7} & \mbox{C08F} & \mbox{C10L} & \mbox{C10M} \end{array}$

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

PAJ, WPI Data, EPO-Internal

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Y	US 5 408 018 A (H. P. RATH) 18 April 1995 (1995-04-18) cited in the application column 1, line 7 -column 2, line 10 column 2, line 44 -column 3, line 18 column 8, line 53 -column 10, line 48 column 10, line 66 -column 11, line 5; claims 1-12	1-88
Y	US 5 448 000 A (P. GULLAPALLI) 5 September 1995 (1995-09-05) the whole document	1-88
Y	EP 0 630 917 A (EXXON RESEARCH AND ENGINEERING COMPANY) 28 December 1994 (1994-12-28) the whole document	1-88

Further documents are listed in the continuation of box C.	Patent tamily members are listed in annex.
A document defining the general state of the art which is not considered to be of particular relevance *E* earlier document but published on or after the international filing date *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) *O* document referring to an oral disclosure, use, exhibition or other means *P* document published prior to the international filing date but later than the priority date claimed	 *T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention. *X* document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone. *Y* document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art. *3* document member of the same patent family
Date of the actual completion of the international search 9 October 2001	Date of mailing of the international search report 22/10/2001
Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nt,	Authorized officer Permentier, W
Fax: (+31-70) 340-3016	Termenerer, n





Interns (at Application No PCT/US 01/08345

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4	EP 0 641 810 A (BASF AG) 8 March 1995 (1995-03-08) claims 1-13		1	
A	EP 0 217 618 A (EXXON RESEARCH AND ENGINEERING COMPANY) 8 April 1987 (1987-04-08) claims 1-17		1	



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